PATENT SPECIFICATION

(21) Application No. 35611/74 (22) Filed 13 Aug. 1974

(31) Convention Application No.

91 664/73 (32) Filed 17 Aug. 1973 in

(33) Japan (JA)

(44) Complete Specification published 8 Sept. 1976

(51) INT. CL. CO8J 5/22

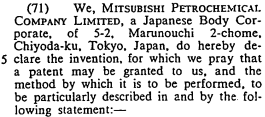
(52) Index at acceptance

C3P 7DIC 7D2A1 7D2A2B 7K7 E2 PC13A PC13B PC13C PC17 PC20D2 T1X T2D

B5A 1G10 1U2AX 2A3 2D1X

C3B 1C12 1C24 1C33A 1C8P 1D2C 1L4E 1L6D





O This invention relates to a process for the production of an ion exchange membrane, generally referred to herein as a "heterogeneous membrane", in which a finely divided ion exchange resin is dispersed in a

15 thermoplastic resin matrix.

In general, press forming, calender forming and extrustion forming are used as methods of forming sheets and films and, in particular, the method of extrusion forming is followed when compared to the other

20 ing is favoured when compared to the other forming methods. The methods of press forming and calender forming can be carried out without trouble on a small scale or laboratory scale, but, in the case of the mass

25 production of large size sheets, it is difficult to control the thickness of the sheet accurately; thus requiring expensive apparatus as well as a high degree of skill. Sheets of even thickness can, by contrast, be easily
30 produced on a large scale or a commercial scale by the method of extrusion forming.

However, it has hitherto been considered impossible to apply the extrusion forming method of the prior art to the production of

35 an ion exchange membrane. The ion exchange resin is so hygroscopic that the water content can amount to from 4 to 20% when allowed to stand in the air, and, therefore, it produces a large amount of gas during ex-

40 trusion inside an extruder. Even if an extruder of the "bent type" is used, extrusion can not be practically carried out because of the large amount of gas produced. Furthermore, an ion exchange resin, being

45 crosslinked, is difficult to melt in an ex-

truder, so that many difficulties arise. These difficulties are the lowering of extrusion capacity, the increase of the back pressure and increased fluciation of the extrusion quantity due to low flow property of the 50 resin; in a manner analogous to the extrusion forming of a resin composition having a high filler content. In order to enhance the electrical conductivity of the resin, which is an important electrochemical property for 55 an ion exchange membrane, it is desirable to increase the quantity of an ion exchange resin being blended; but the increase of the quantity of the ion exchange resin results inevitably in difficulties due to the increas- 60 ing of the hygroscopic property and lowering of the flow property, as described above.

Efforts have been made to overcome these difficulties and consequently a way has been 65 found to produce an ion exchange membrane on a commercial scale by extrusion. The present invention is based on this discovery.

The above mentioned difficulties can be 70 overcome by a process for the production of an ion exchange membrane consisting of an ion exchange resin finely and heterogeneously dispersed in a thermoplastic resin.

Accordingly the present invention pro- 75 vides a process for the production of an ion exchange membrane comprising a finely divided ion exchange resin heterogeneously dispersed in a thermoplastic resin, which process comprises mixing a thermoplastic resin 80 with substantially dry ion exchange resin powder in the proportions of from 2:8 to 8: 2 parts by weight respectively, preparing pellets from the mixture, adding a lubricant to the mixture, before or after the pelletizing, 85 selected from a metallic soap, a fatty acid amide or a fatty acid ester in a proportion of from 0.1 to 10 parts by weight per 100 parts. by weight of the sum of the weights of the thermoplastic resin and the ion exchange 90



105

resin; and then subjecting the pellets to extrusion and to pressing using at least one set of rolls maintained at a temperature of from 50 to 150°C to form the membrane.

In accordance with the process of the present invention, the extrusion forming or moulding can be carried out on a commercial scale, since the ion exchange resin is pelletized and sealed in a thermoplastic 10 resin, resulting in a marked decrease of the hygroscopic property of the resin. In addition, a lubricant is used which is capable of improving the lubricating property at the boundary of the resin, thus improving the

15 flow property of the resin as well as increasing the efficiency of the extrusion process.

The process of the present invention will now be described in more detail.

The basic starting materials of the process according to the present invention are an ion exchange resin, a thermoplastic resin, a lubricant and, if necessary, an additive such as a stabilizer.

The ion exchange resin used may be a cation exchange resin suitable for the production of cation exchange membranes, an anion exchange resin for the production of anion exchange membranes, or a mixture of a cation exchange resin and an anion exchange resin, for the production of the ion exchange resin, for the production of the ion exchange membranes for special uses. The cation exchange resins used may be formed from synthetic resins whose base polymers consist of polystyrene, poly (styrene/butadiene), polyvinyt chloride or a phenol-aldehyde resin, and whose ion exchange groups are sulfonic acid, phosphonic acid, phosphinic

40 acid or carboxylic acid groups. The anionic exchange resins used may be formed from synthetic resins whose base polymers consist of an epoxly resin, polyvinyl chloride, polystyrene, or a phenol-aldehyde resin and

45 whose exchange groups are primary to quaternary amino groups. Above all, it is desirable to use an ion exchange resin having a degree of swelling of 0.5 or more, preferably 3.0 or more, and the resin should 50 be stable at the melting point of a thermo-

50 be stable at the melting point of a thermoplastic resin. By the term "degree of swelling" as used herein is meant the ratio of the volume of the resin in a swollen state to that in a shrunken state during use there-

55 of, as used commonly in the field of ion exchange resins. The ion exchange resin used should have a sufficiently fine particle size and should preferably be a powder of a particle size small than 100 Tyler mesh.

60 The choice of the thermoplastic resin used is not particularly critical. Any thermoplastic resin which is comprised of polyethylene, polypropylene, an ethylene-vinyl acetate copolymer; polyvinyl chloride, poly-65 styrene or a polyamide may be used, but in

particular, the use of crystalline polyolefins and olefin copolymers such as low density polyethylene, high density polyethylene, polypropylene and the ethylene-propylene copolymers is desirable.

The above mentioned thermoplastic resin and ion exchange fine powder are mixed in a proportoin of from 2:8 to 8:2, preferably from 3:7 to 6:4. If the proportion of the ion exchange resin used is less 75 than that given in this range, then the ion exchanging property of the membrane cannot be obtained; while if greater, the formation of a coherent membrane will be found to be impossible.

In a preferred embodiment of this invention, an ion exchange resin, a thermoplastic resin and the required additives are mixed or blended while exhausting the generated gas by means of an ordinary melting and 85 blending apparatus such as a kneader or a mixing roll. The blend is then formed in a sheet using rolls and then cut to prepare a pellet-shaped raw material suitable for extrusion. The resulting pellets can be 90 handled as a raw material for ordinary extrusion moulding, since the water content thereof is in general reduced to about 2% and the rate at which moisture is absorbed is much lower than in the case of an ion 95 exchange resin alone. This mixing step is generally carried out using the following operations:

(a) effective drying of a fine powder of ion exchange resin,

(b) dispersion of the fine resin powder,
(c) sealing of the fine resin powder in a thermoplastic resin, thus rendering the ion exchange resin less hygroscopic.

In general, ion exchange resins have a very poor resistance to heat, but, during the mixing and extrusion moulding steps of the present invention, little deterioration takes place upon heating at a temperature of up 110 to 240°C for up to 45 minutes.

It has also been found that the addition of a particular lubricant is markedly effective for improving the flow property and workability of such a resin. The effects are as 115 follows:

(a) an increase of the rate of extrusion,
(b) a decrease of the fluctuation of the resultant membrane thickness due to changes in the quantity extruded,

(c) a suppression of the crack formation that often occurs during thin membrane moulding.

This lubricant is either a metallic soap, a fatty acid amide or a fatty acid ester. The 125 metallic soap used may be a metal salt of a fatty acid having from 10 to 22 carbon atoms, such for example as the calcium, vanadium, cadmium or magnesium salt of stearic acid, palmitic acid or oleic acid. The 130

3

olyolefins w density yethylene, propylene ısı esin mixed in : 2, pree propored is less 75 n the ion rane canhe formabe found his invenmoplastic are mixed generated ilting and 85 ader or a med in a o prepare itable for can be 90 linary exer content about 2% absorbed of an ion 95 ig step is following powder of 100 /der. waer in a rendering :ss hygro-105 is have a during the eps of the tion takes ure of up 110 addition y effective and workcts are as 115 extrusion, on of the ess due to truded. 120 formation hin memallic soap, ester. The 125 tal salt of 22 carbon : calcium,

m salt of

acid. The 130

fatty acide amide used may be a compound represented by the general formula R-CO-NH₂ in which R represents a saturated or unsaturated hydrocarbon group having 5 from 1 to 21 carbon atoms, or a bis-fatty acid amide represented by the general formula: R₁—CO—NH—R₂—NH—CO—R₃ in which R₁ and R₂ each individually represent a saturated and unsaturated hydrocar-10 bon group having from 9 to 21 carbon atoms and R₂ represents a saturated hydrocarbon group having from 1 to 5 carbon atoms. The fatty acid ester used may be a compound represented by the general formula: 15 R₁(OH)_n—OOC—R₂ in which R₁ represents a straight chain or cyclic saturated or unsaturated hydrocarbon group having from 1 to 22 carbon atoms and R₂ represents a saturated or unsaturated hydrocarbon group 20 having from 9 to 21 carbon atoms, and n represents 0 or an integer of 1 or more. Examples of these compounds are oleic acid amide, stearic acid ethylene bisamide, stearic acid monoglyceride, behenic acid mono-25 glyceride, ricinoleic acid monoglyceride, oleic acid monoglyceride, lauric acid sorbitan ester, palmitic acid sorbitan ester and oleic acid sorbitan ester. Sorbitan esters of saturated or unsaturated fatty acids are par-30 ticularly preferable. It is assumed that since each of the lubricants of a metallic soap, a fatty acid amide or a fatty acid ester type has an alkyl attracting portion and polar attracting por-35 tion, an ion exchange resin having a strong polarity can uniformly be dispersed in a thermoplastic resin and, simultaneously, the lubricating property at the boundary of the resin can be improved to increase 40 the flow property of the resin. The electrochemical property of an ion exchange membrane is not lowered even by the addition of such a lubricant. The lubricant is added in a proportion 45 of from 0.1 to 10 parts by weight, preferably from 2.0 to 4.0 parts by weight per 100 parts by weight of the sum of the weights of the ion exchange resin and thermoplastic resin. If the lubricant is added 50 in a proportion of less than 0.1 part by weight, no effect is observed by the addition thereof and a uniform membrane is difficult to mould by extrusion; while if

more than 10 parts by weight, the lubri-

is sometimes lowered. The addition of the

lubricant is ordinarily carried out by add-

step or by adding it to the mixture before

or after the peletizing step. In view of the

ease of operation, the former addition

The thus prepared pellets are fed to an

60 ing it to the raw material before the mixing

method is preferred.

55 cating effect is not significantly increased further and, in addition, the electro-chemical exchange property of the resin

3 extruder equipped with an ordinary T die to form a membrane which is then subjected to a pressing treatment using rolls such as calendering rolls, followed by rolling-up of the finished membrane. This 70 pressing treatment is carried out for the purpose of cooling and solidifying the membrane and preventing the membrane from shedding some of the ion exchange resin particles during cooling and solidify- 75 ing and from thus forming pinholes; as well as for giving the surface a smooth and lustrous appearance. The surface temperature of the roll should be kept within a range of from 50 to 150°C, since at below 80 50°C, the surface of the membrane roughens markedly and, in an extreme case, cracks or pinholes are formed; while at above 150°C, the cooling of the membrane is insufficient, thus resulting in 85 adhesion of the membrane to the roll surface and hence deformation of the membrane. A temperature of from 70 to 130°C is particularly preferably. The pressing treatment is preferably carried out in a 90 multistage process using from two to five pairs of rolls. The resultant membrane is slightly foamed, but it can be used as an ion exchange membrane as it is. In order to 95 obtain better performance of the ion exchange membrane, the membranes may be subjected to an after-treatment step comprising immersion of the membrane in either a swelling liquid such as an acidic 100 alkaline solution containing, for example, from 1 to 20% by weight of hydrochloric acid or caustic soda at a temperature of 60°C or higher, or hot water at a temperature of 80°C or higher 105 for several minutes or more. The following Examples further illustrate the present invention. The mesh standard referred to in these Examples is the Tyler mesh standard. Comparative Example 1 90 parts by weight of styrene monomer and 10 parts by weight of divinylbenzene monomer were copolymerized to give a bridged polystyrene in the form of beads 115 about 1 mm in diameter. These were then sulfonated in a known manner to obtain a cation exchange resin, and then pulverized. The portion of the cation exchange resin passing through a sieve of 100 mesh was 120

used as a raw material, which had a water content of 15% by weight. To 60 parts by weight (on dry base) of this ion exchange resin were added 3 parts by weight of powdered sorbitan stearate as a lubricant 125 followed by mixing for about 10 minutes in a mixer of the high speed revolving blade type. Then 40 parts by weight of powdered polypropylene having a melt index of 5 g/10 min at 230°C was added 130

115

thereto and mixed for 10 minutes. The mixed raw material was subjected to extrusion moulding using an ordinary extruder but this extrusion could not be 5 carried out without danger because of small explosions occurring in the cylinder. Thus the extrusion was accomplished using an extruder of bent type (40 φ , L/D = 27) equipped with an ordinary T die. The 10 optimum conditions were found and used. These were a temperature range of from 155 to 165°C under the hopper, from 180 to 205°C at the bent portion, from 200 to 230°C at the second compression portion 15 and metering portion and from 200 to 230°C at the die portion. Even under the best condition, the quantity of extrusion reached only 6 kg/hr with a screw revolving at 60 rpm and the fluctuation of the rate 20 of extrusion was so large (± 15% width) that an even, continuing membrane could not be obtained. Furthermore, a vigorous venting of the resin powder took place from the vent hole at the same time.

Example 1 In a manner analogous to Comparative Example 1, 60 parts by weight of the ion exchange resin passing through 100 mesh, 3 parts by weight of sorbitan stearate and 30 40 parts by weight of powdered polypropylene were mixed, kneaded at 185°C for about 30 minutes by means of an ordinary mixing roll and then formed into pellets by the sheet cutting method. The 35 resulting pellets were fed to an extruder provided with an ordinary T die (45 ϕ , $\hat{L}/D = 25$, extruded in the form of a membrane and pressed by passing through three stage rolls heated to 120°C to obtain 40 in a stable manner a uniform product 0.25 mm in thickness. The rate of extrusion was 11 kg/hr with a screw revolving it 60 rpm and the fluctuation thereof was only up to : ± 2%.

Example 2 In a manner analogous to Example 1, 60 parts by weight of the ion exchange resin passing through 100 mesh, 3 parts by weight of ethylene-N.N'-methylene-bis-50 stearamide as a lubricant and 40 parts by weight of powdered polypropylene were mixed and formed into pellets, and then extruded by means of the same apparatus as that of Example 1 thus obtaining a 55 uniform product 0.35 mm in thickness in a stable manner. The rate of extrusion reached 9.8 kg/hr with a screw revolving it 60 rpm and with a fluctuation of up to ± 2%.

Comparative Example 2 For comparison with Examples 1 and 2, pellets prepared from 60 parts by weight of the ion exchange resin and 40 parts by weight of polypropylene without adding a 65 lubricant were moulded by the same appa-

ratus as that of Example 1 or 2. However, the fluctuation of the rate of extrusion reached 1± 10% or more and cracks were produced from a lower strength area which had been just extruded from the die lips 70 and had not cooled and solidified sufficiently. Thus no product was given. The rate of extrusion was 9.1 kg/hr. When the die temperature was raised to 150°C, some improvement was found with regard to the 75 formation of cracks but the problem could not be solved fundamentally.

Comparative Example 3 A membrane was obtained by extruding the same raw material in the same manner 80 as in Example 1 and then the membrane was passed through a three stage roll of iron plated with hard chromium to obtain a product of 0.41 mm in thickness. The surface of the membrane lacked smooth- 85 ness and lustre and had irregular waved patterns and pinholes on very uneven areas.

Example 3 When the membrane-shaped product of Example 1 was treated with hot water for 90 I hour and then washed with water, the transport number of the membrane was 0.95 in a 0.5 N NaCl solution and the specific resistance was $180\Omega cm$.

95 Example 4 A fine powder of an anion exchange resin consisting of an epoxy resin as base polymer having amino groups as the exchange groups and passing through 100 mesh was prepared in a known manner. 60 parts by 100 weight of the anion exchange resin, 3 parts by weight of sorbitan stearate as a lubricant and 40 parts by weight of powdered polypropylene were mixed and moulded in a manner analogous to Example 1; thus 105 obtaining a uniform product 0.25 mm in thickness in a stable manner. The rate of extrusion was 12.3 kg/hr with a screw revolving it 60 rpm and with an extrusion fluctuation of ±3%. The resulting mem- 110 brane had a transport number of 0.93 in a 0.5 N NaCl solution and a specific resistance of 93Ωcm after being treated with hot water for 1 hour and washed with water.

WHAT WE CLAIM IS: -1. A process for the production of an ion exchange membrane comprising a resin exchange ion divided finely heterogeneously dispersed in a thermo- 120 plastic resin, which process comprises mixing a thermoplastic resin with substantially dry ion exchange resin powder in the proportions of from 2:8 to 8:2 parts by weight respectively, pre-paring pellets from the mixture, adding a lubricant to the mixture before or after the pelletizing, selected from a metallic soap, a fatty acid amide or a fatty acid ester in a proportion of from 130

wever, trusion s were variety in the some to the 75 could

truding nanner 80 nbrane roll of obtain s. The mooth-waved 1 areas.

luct of ter for 90 er, the ne was nd the

95 ze resin e poly-:change st ras oy 100 3 parts *ibricant* d polyd in a ; thus 105 mm in rate of screw trusion mem- 110 0.93 in specific treated ed with 115

of an sing a resin thermo- 120 es mix-1 sub-powder 1:8 to y, pre-125; addre be-selected nide or y' om 130

0.1 to 10 parts by weight per 100 parts by weight of the sum of the weights of the thermoplastic resin and the ion exchange resin; and then subjecting the pellets to extrusion and to pressing using at least one set of rolls maintained at a temperature of from 50 to 150°C to form the mebrane.

A process as claimed in claim 1 wherein the thermoplastic resin and the ion 10 exchange resin are mixed in the proportion of from 3: 7 to 6: 4 parts by weight respectively.

3. A process as claimed in claim 1 or claim 2 wherein the lubricant is added to 5 the resin mixture in the proportion of from 2.0 to 4.0 parts by weight of lubricant to 100 parts by weight of the sum of the weights of the thermoplastic resin and the ion exchange resin.

4. A process as claimed in any one of claims 1 to 3 wherein the pressing step is carried out using from two to five pairs of rolls heated to a temperature of from 70 to 130°C.

25 5. A process as claimed in any one of the preceding claims wherein the lubricant is a sorbitan ester of a saturated or unsaturated fatty acid.

6. A process as claimed in any one of 30 the preceding claims wherein the thermoplastic resin is a crystalline olefin polymer or olefin copolymer.

7. A process as claimed in any one of the preceding claims wherein a stabilizer is 35 additionally present in the resin mixture.

8. A process as claimed in any one of the preceding claims wherein the ion exchange resin has a degree of swelling of greater than or equal to 0.5.

40 9. A process as claimed in claim 8 wherein the ion exchange resin has a degree of swelling of greater than or equal to 3.0.

10. A process as claimed in any one of the preceding claims wherein the ion ex- 45 change resin has a particle size of less than 100 Tyler mesh.

11. A process as claimed in any one of the preceding claims wherein the pressing step is followed by an after-treatment step 50 which comprises the immersion of the membrane in a swelling liquid.

12. A process as claimed in claim 11 wherein the swelling liquid is an acidic solution having a temperature of 60°C or 55 higher.

13. A process as claimed in claim 12 wherein the acidic solution is an aqueous solution containing from 1 to 20 percent by weight of hydrochloric acid.

14. A process as claimed in claim 11 wherein the swelling liquid is an alkaline solution having a temperature of 60°C or higher.

15. A process as claimed in claim 65 14 wherein the alkaline solution is an aqueous solution containing from 1 to 20 per cent by weight of sodium hydroxide.

16. A process as claimed in claim 11 wherein the swelling liquid is water having 70 a temperature of 80°C or higher.

17. A process as claimed in claim 1 substantially as hereinbefore described in any one of the Examples.

18. An ion exchange membrane when 75 produced by a process as claimed in any one of the preceding claims.

MITSUBISHI PETROCHEMICAL COMPANY LIMITED,

Per

BOULT, WADE & TENNANT, 34 Cursitor Street, London, EC4 1PQ. Chartered Patent Agents.

Printed for Her Majesty's Stationery Office by The Tweeddale Press Ltd., Berwick-upon-Tweed, 1976. Published at the Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.